(51) International Patent Classification 7:		(11) International Publication Number:	WO 00/22189
C23F 11/10, 11/18, C09K 5/20	A1	(43) International Publication Date:	20 April 2000 (20.04.00)
(21) International Application Number: PCT/IB9 (22) International Filing Date: 11 October 1999 (1 (30) Priority Data: 98308380.9 14 October 1998 (14.10.98) (71) Applicant (for all designated States except US): TEXA VELOPMENT CORPORATION [US/US]; 2000 ester Avenue, White Plains, NY 10650 (US).	1.10.9 I	BR, BY, CA, CH, SN, CU, C GD, GE, GH, GM, HR, HU, KP, KR, KZ, LC, LK, LR, LS, MN, MW, MX, NO, NZ, PL SI, SK, SL, TJ, TM, TR, TT, ZA, ZW, ARIPO patent (GH, SZ, TZ, UG, ZW), Eurasian pa MD, RU, TJ, TM), European p DK, ES, FI, FR, GB, GR, IE	Z, DE, DK, EE, ES, FI, GB ID, IL, IN, IS, JP, KE, KG LT, LU, LV, MD, MG, MK PT, RO, RU, SD, SE, SG UA, UG, US, UZ, VN, YU GM, KE, LS, MW, SD, SL tlent (AM, AZ, BY, KG, KZ totatent (AT, BE, CH, CY, DE IT, LU, MC, NL, PT, SE
 (72) Inventors; and (75) Inventors/Applicants (for US only): MAES, Jea [BE/BE]; Burgemeester De Guchteneerelaan 36, Merelbeke (BE). LIEVENS, Serge [BE/BE]; Sche pontweg 22, B-9820 Merelbeke (BE). (74) Agents: GREEN, Mark, Charles et al.; Urquhart-I Lord, 91 Wimpole Street, London W1M 8AH (GB 	B-983 ellebell Dykes	20 With international search repo	rt.

(54) Title: CORROSION INHIBITORS AND SYNERGISTIC INHIBITOR COMBINATIONS FOR THE PROTECTION OF LIGHT METALS IN HEAT-TRANSFER FLUIDS AND ENGINE COOLANTS

(57) Abstract

Corrosion protection of magnesium and aluminium alloys in engine coolants and heat-exchange fluids is achieved by the use of a select group of aliphatic and aromatic carboxylate acids or the alkaline metal, ammonium or amine salts thereof in combination with a fluoride and/or a fluorocarboxylic acid or salt thereof. These compositions have been found to significantly improve the high temperature magnesium corrosion protection properties of coolants, and are of use in automobile engine coolant systems.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL AM AT AU AZ BA BB BE BF BG BJ BR CA CF CG CM CN CU CZ DE DE EE	Albania Armenia Australia Australia Azerbaijan Bosnia and Herzegovina Barbados Belgium Burkina Faso Bulgaria Benin Brazil Belarus Canada Central African Republic Congo Switzerland Côte d'Ivoire Cameroon China Cuba Czech Republic Germany Denmark Esionia	ES FI FR GA GB GE GN GR HU IE IL IS IT JP KE KG KP KR LC LL LK LR	Spain Finland France Gabon United Kingdom 4 Georgia Ghana Guinea Greece Hungary Ireland Israel Iceland Italy Japan Kenya Kyrgyzstan Democratic People's Republic of Korea Republic of Korea Kazakstan Saint Lucia Liechtenstein Sri Lanka Liberia	LS LT LU LV MC MD MG MK ML MN MR MW MX NE NL NO NZ PL PT RO RU SD SE SG	Lesotho Lithuania Luxembourg Latvia Monaco Republic of Moldova Madagascar The former Yugoslav Republic of Macedonia Mali Mongolia Mauritania Malawi Mexico Niger Netherlands Norway New Zealand Poland Portugal Romania Russian Federation Sudan Sweden Singapore	SI SK SN SZ TD TG TJ TM TR TT UA UG US UZ VN YU ZW	Slovenia Slovakia Senegal Swaziland Chad Togo Tajikistan Turkmenistan Turkey Trinidad and Tobago Ukraine Uganda United States of America Uzbekistan Viet Nam Yugoslavia Zimbabwe
---	--	---	---	---	---	--	--

15

20

25

CORROSION INHIBITORS AND SYNERGISTIC INHIBITOR COMBINATIONS
FOR THE PROTECTION OF LIGHT METALS IN HEAT-TRANSFER FLUIDS
AND ENGINE COOLANTS.

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to compositions which assist in the corrosion protection of magnesium, magnesium-aluminum, aluminum-magnesium and aluminum alloys in engine coolants and heat-exchange fluids. Aliphatic mono- or di-basic acids or aromatic carboxylate acids, or the alkali metal, ammonium or amine salts thereof, have been found to provide corrosion protection to magnesium, in addition to the protection that these acids already provide to other metals, such as aluminum, iron, copper and solder, when combined with a fluoride or a fluorocarboxylic acid or a salt thereof. Specific combinations of such acids or salts with fluoro compounds have been found to provide synergistic corrosion protection for magnesium. optional addition of a hydrocarbyl triazole and/or a thiazole to these combinations provides improved corrosion protection, particularly to copper alloys but also to other metals such as The presence of the fluoride fluorocarboxylate has been found to significantly improve the high temperature magnesium corrosion protection properties for individual carboxylates, for combinations of carboxylates, and for combinations of carboxylates and hydrocarbyl triazoles and/or thiazoles.

Automobile engine cooling systems contain a variety of metals, including copper, solder, brass, steel, cast iron, aluminium,

10

15

20

25

magnesium, and their alloys. The possibility of corrosive attack on such metals is high, due to the presence of various ions as well as the high temperatures, pressures, and flow rates found in such cooling systems. The presence of corrosion products within the cooling system can interfere with heat transfer from the engine combustion chambers, which may subsequently cause engine overheating and engine component failure due to excess metal temperatures. See generally Fay, "Antifreezes and Deicing Fluids", Kirk-Othmer Encyclopaedia of Chemical Technology (1978) ed, vol 3, pp 79-95. therefore be generally advantageous if the formation of corrosion products within automobile cooling systems could be It is one object of the instant controlled or eliminated. invention to provide a corrosion inhibitor useful in the prevention and control of corrosion in automobile engine cooling systems containing various metals, particularly magnesium.

The trend towards improved fuel economy for automobiles has led to the increased use of lightweight materials such as aluminium and magnesium alloys for engine and cooling system components. However, it has been found that pitting and crevice corrosion are particularly prevalent in aluminium and magnesium-containing cooling systems. Pitting of thin-walled automobile radiator tubes may lead to tube perforation. Crevice corrosion at cylinder head packings or coolant hose connections may also occur. Both types of corrosion may lead to eventual coolant loss, with subsequent engine overheating and component failure. Other forms of localized corrosion such as deposit attack from deposition of corrosion products may also result.

.5

10

15

20

25

Many conventional corrosion inhibitor additives used in automobile cooling systems do not provide adequate protection against the pitting, crevice, and deposit attack corrosion phenomena found with magnesium, aluminium and various other metal alloys. It would therefore be particularly advantageous if such localized corrosion phenomena could be controlled or eliminated. It is another object of the instant invention to provide a corrosion inhibitor for use in automobile cooling systems which prevents or controls localized corrosion of magnesium.

All corrosion inhibitors employed in automobile antifreeze/coolant formulations are gradually depleted by use and the build-up of corrosion products in the cooling system. It would thus be advantageous if the build-up of corrosion products within the system and subsequent corrosion inhibitor depletion or degradation could be controlled or eliminated. It is a further object of the instant invention to provide a corrosion inhibitor which is less prone to depletion or degradation than traditional corrosion inhibitors used in antifreeze/coolant formulations.

2. Description of Related Information

Organic Acid Technology (OAT) coolants and heat exchange fluids have been introduced, providing improved corrosion protection and having a long life. OAT corrosion-inhibitor packages in aqueous and glycol concentrates are used in automotive, heavy duty, marine and industrial applications. OAT corrosion-inhibitors are also used in secondary cooling systems and in a variety of industrial heat exchange fluids. Several U.S. and

10

15

20

25

WO 00/22189 PCT/IB99/01659

foreign patent references disclose the use of carboxylic acids, or the salts of such acids as corrosion inhibitors in antifreeze/coolant and heat-exchange fluid compositions. These compositions are optimized for the protection of aluminum and other materials currently used in the above applications.

Various corrosion inhibitors have been added to heat transfer fluids to reduce corrosion of metallic systems. For example, discloses non-silicate Patent No 4,587,028 (Darden) antifreeze formulations containing alkali metal salts of dicarboxylic acid and nitrate. Additional benzoic acid, ingredients including alkali metal hydroxides, alkali metal nitrates and aromatic triazoles, such as tolyltriazole or benzotriazole are preferably provided. US Patent No 4,647,392 (Darden et al) discloses corrosion inhibitors using aliphatic monobasic acids or salts, hydrocarbyl dibasic acids or salts and hydrocarbonyl triazole. US Patent No 4,657,689 (Darden) discloses corrosion inhibitors containing aliphatic monobasic acids or salts, hydrocarbyl dibasic acids or salts, hydrocarbyl azoles and specific hydrocarbyl alkali metal sulfonates. Patent No 5,085,791 (Burns) discloses antifreeze compositions containing cyclohexane acid corrosion inhibitor alone or in combination with other corrosion inhibitors, particularly sebacic acid and tolyltriazole. The cyclohexane acid includes cyclohexyl carboxylic (formic) acid, cyclohexyl acetic acid and cyclohexyl propionic acid. The cyclohexane acid is targeted to inhibit lead solder and/or aluminium corrosion. US Patent No 4,105,405 (Wehle et al) discloses the use of cyclohexane hexacarboxylic acid corrosion inhibitors.

10

15

20

25

JP-A-08 020763 (Seiken Kogaku Kogyo KK) describes a fluid for use as a coolant in internal combustion engines comprising glycol, water, a magnesium compound, an alkyl benzoic acid (e.g. p-tert-butylbenzoic acid) or a salt, a do-decane diacid or a salt and a triazole or thiazole.

JP-A-08 085782 (Nippon Chem Kogyo KK) describes a glycol based antifreeze composition containing a dodecanedioic acid or a salt, p-tert-butylbenzoic acid or a salt and a triazole, together with a silicate, molybdate, benzoate or thiazole. The composition is free from amines, phosphates, borates and nitrites.

Engine manufacturers are now evaluating the use of magnesium as a material for engine and heat-transfer systems. Traditional inhibitor packages do not give adequate corrosion protection to magnesium components. In general, currently used OAT coolants are only mildly aggressive, but the protection levels for magnesium are not sufficient, particularly at the high temperatures found in working engines. Previous research has alkylbenzoic acid combination of an a that indicated (4-tert-butylbenzoic acid), an aliphatic monoacid (octanoic acid) and a hydrocarbyl triazole (tolyltriazole), provides improved corrosion protection for magnesium, in comparison with traditional and OAT coolant formulations (Table I in US Patent 4,851,145 (van Neste)). EP-A-0229254 (Asahi Glass Co Ltd) describes an electrolytic capacitor comprising a capacitor element and an electrolyte impregnated into the element, wherein the electrolyle contains a fluorocarboxylic acid or salt dissolved in an organic solvent.

10

15

20

There is need for a coolant system which provides high levels of corrosion protection to magnesium components.

SUMMARY OF THE INVENTION

The present invention relates to an antifreeze concentrate composition comprising one or more inhibitors selected from alkylbenzoic acid, C_5 - C_{15} mono- or di-basic acid or salts thereof, together with a fluoride and/or a fluorocarboxylic acid or salt thereof.

Amongst the aromatic carboxylates, the group of alkylbenzoic acids of general formula

where R is a C_1 - C_8 alkyl radical or an element of group 7, e.g. F, are preferred. 4-Tert-butylbenzoic acid (hereafter referred to as PTBBA), is the most preferred alkylbenzoic acid and 4-fluoro benzoic acid is the most preferred compound when R is an element of group 7. Alkali metal, ammonium or amine salts of the alkylbenzoic acid may be used. The most preferred alkali metal salts are potassium and sodium.

Aliphatic carboxylate acids that provide corrosion protection may be any C_5 - C_{15} aliphatic monobasic acid or dibasic acid or the alkali metal, ammonium or amine salt of said acids.

15

20

25

Heptanoic acid, octanoic acid, and nonanoic acid (or isomers thereof), and mixtures thereof are the preferred monobasic acids. Decanoic acid and undecanoic acid provide good protection but solubility of the higher alkyl acids in water is limited. Of the dibasic acids, dodecanedioic acid provides reasonably good magnesium protection. Of the naphtylcarboxylic acids, 1-naphtylcarboxylic acid is the preferred carboxylic acid.

It has been found that the combination of one or more of the above described acids, together with the fluoro compound, gives a synergetic effect for improved magnesium protection. The combination of PTBBA and octanoic acid is especially preferred. Nonanoic and heptanoic acid are good alternatives for octanoic acid. Compositions formulated with no hydrocarbyl triazole give good magnesium protection, or hydrocarbyl triazole at a level below 0.2 wt%. The addition of a hydrocarbyl triazole to these combinations provides additional copper protection, as expected. Improved corrosion protection properties were found for the other metals, especially for aluminum.

It has been found that the addition of a fluoride or a fluoro carboxylate to an aliphatic carboxylic acid or to an alkylbenzoic acid or to the preferred synergistic combinations of alkylbenzoic acids and the preferred monocarboxylic acids, significantly improves the high temperature corrosion protection properties of the formulation. Most particularly, such systems give improved protection against magnesium corrosion.

10

15 .

20

DETAILED DESCRIPTION OF THE INVENTION

The present invention firstly identifies a group of specific aliphatic and aromatic carboxylates that provide improved corrosion protection, when used with a fluoride and/or a fluorocarboxylic acid or salt thereof.

Amongst the aromatic carboxylates are preferred the group of alkylbenzoic acids of general formula

where R is a C_1 - C_6 alkyl radical. In this invention, 4-tert-butylbenzoic acid (PTBBA) is preferred. Any alkali metal, ammonium or amine salts of the alkylbenzoic acid may be used to form the alkylbenzoic acid salt; however, alkali metals are preferred. The most preferred alkali metal salts are potassium and sodium. Of the naphtylcarboxylic acids, 1-naphtylcarboxylic acid is the preferred carboxylic acid.

Aliphatic carboxylate acids that provide corrosion protection may be any C_5 - C_{15} aliphatic monobasic acid or dibasic acid or the alkali metal, ammonium or amine salt of said acids. This would include one or more of the following acids or isomers thereof: heptanoic acid, octanoic acid, and nonanoic acid, and mixtures thereof are the preferred monobasic acids. Decanoic acid and undecanoic acid provide good protection but solubility of the higher alkyl acids in water is limited. Of the dibasic

acids, dodecanedioic acid provides reasonably good magnesium protection.

Combinations of one or multiples of the above mentioned aromatic carboxylates with one or more aliphatic carboxylates, together with the fluoro compound, give a synergistic effect for the protection of magnesium alloys. When using a hydrocarbyl triazole component such as an aromatic triazole or an alkyl substituted aromatic triazole such as benzotriazole or tolyltriazole, increased protection of copper is achieved. Improved protection of other metals, particularly aluminum is also observed. For magnesium protection alone the use of a hydrocarbyl triazole is optional.

10

15

20

25

inhibitor which now provides superior The additional synergistic corrosion protection, especially for magnesium alloys, especially at elevated temperatures, in combination with one carboxylic acid or with a synergetic mixtures of the combination of aliphatic multiple acids such as carboxylates with the alkylbenzoic acids, such as mentioned above, is fluoride. Used alone, fluoride provides a very low It is an important aspect of this level of protection. invention that fluor compounds such as fluorides and/or fluoro carboxylates, in combination with aliphatic carboxylates or alkylbenzoic acids give a synergistic effect to significantly magnesium at elevated reduce corrosion, especially of The fluoride can be introduced formulation as hydrogen fluoride and/or a fluoro carboxylic acid or the alkali metal, ammonium or amine soluble salt of the mentioned acid.

The combination of the corrosion inhibitors of the instant invention will most typically be employed in antifreeze formulations as a coolant for internal combustion engines. Other applications may include hydraulic fluids, aqueous cutting oils, paints, soluble oils, metal cutting fluids, aircraft deicers, and greases. In these applications the aliphatic acids and alkylbenzoates may be formed with metal hydroxides including sodium, potassium, lithium and barium.

5

10

15

20

25

In a preferred embodiment of the instant invention, the above described corrosion inhibitors may be employed in a mixture with a liquid alcohol freezing point depressant to form a novel antifreeze/coolant concentrate composition for use in the cooling systems of internal combustion engines. The coolant concentrate composition preferably comprises: from 80-99 weight percent of a water soluble liquid alcohol freezing point depressant; from 0.01-15, preferably 0.1-5 weight percent of the above described alkylbenzoic acid/salt and/or aliphatic carboxylic acid/salt component and from 0.005-5, preferably 0.01-1 weight percent of hydrogen fluoride and/or a fluoro carboxylate. In addition one of the above mentioned components, hydrocarbyl triazole components can optionally be used at from 0.005-1, preferably 0.1-0.3 weight percent.

The liquid alcohol freezing point depressant component of the above described coolant in the instant invention includes glycols such as ethylene glycol, di-ethylene glycol, propylene glycol, di-propylene glycol and glycol monoethers such as the methyl, ethyl, propyl and butyl ethers of ethylene glycol, di-ethylene glycol, propylene glycol and di-propylene glycol. Ethylene and propylene glycol are particularly preferred as the

10

15

20

25

freezing point depressant component. In the above described coolant concentrate composition of the instant invention, an additional dicarboxylic acid preferably dodecanoic diacid, may be employed at concentrations of 0.01-15.0 weight percent, preferably 0.1-5.0 weight percent. Additional conventional corrosion inhibitors such as alkali metal borates, silicates, benzoates, nitrates, nitrites, molybdates or hydrocarbyl thiazoles may also be employed at concentrations of 0.01-5.0 weight percent.

In another embodiment of the invention, the above described corrosion inhibited coolant concentrate composition is diluted with 10-90 volume percent, preferably 25-75 volume percent of water. In yet another embodiment of the invention, the above described corrosion inhibitor combinations are used in aqueous solutions of alternative freezing point depressants such as organic acid salt solutions. Acetates, formates and proprionates are particularly preferred.

EXAMPLES

The method of the invention will be further illustrated by the following examples, which are not intended to limit the scope of the invention.

Comparative Examples

Example A (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent 2-ethyl hexanoic acid:

Example B (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 0.2 weight percent tolyltriazole.

Example C (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 0.2 weight percent HF (50 weight percent in water).

Example D (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent heptanoic acid.

Example E (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent octanoic acid.

Example F (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent dodecanedioic acid.

Example G (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent sebacic acid.

Example H (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent octanoic acid, 0.2 percent TTZ.

Example I (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent PTBBA, 0.2 percent TTZ.

Example J (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent octanoic acid, 0.2 weight percent TTZ.

Example K (Comparative Example)

An antifreeze formulation was prepared comprising a major 15 amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent octanoic acid, 0.02 weight percent TTZ.

Example L (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent octanoic acid.

PCT/IB99/01659

5

10

15

20

Example M (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent octanoic acid, 0.5 percent 2-ethylhexanoic acid, 0.2 weight percent TTZ.

Example N (Comparative Example)

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent octanoic acid, 0.1 weight percent sodium metasilicate pentahydrate, 0.2 weight percent TTZ

Examples of the Invention

Example 1

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight percent OA, 0.2 weight percent HF (50 weight percent in water), 0.2 weight percent TTZ.

Example 2

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent PTBBA, 0.2 weight percent HF.

Example 3

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent OA, 0.2 weight percent HF.

Example 4

5

15

20

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 3 weight percent heptanoic acid, 0.2 weight percent HF.

Example 5

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5 weight fluorobenzoic acid.

Example 6

An antifreeze formulation was prepared comprising a major amount of ethylene glycol, 1.5 weight percent PTBBA, 1.5.weight percent octanoic acid, 0.2 weight percent sebacic acid, 0.2 weight percent sodium fluoride acid, 0.2 weight percent TTZ.

The additive systems are summarised in Table 1.

It is well known that corrosion resistance of a metal or metal alloy depends upon both the stability of its passivating oxide protective film and its ability to repassivate active corrosion regions on the surface of the metal or alloy. The speed of

corrosion on the other hand is related to the current density. A rapid Cyclic Polarization Scanning (RCP) technique, based on cyclic potentiokinetic polarization and described in the CEBELCOR (Centre Belge d'Etude de la corrosion) publication Rapports Techniques, Vol. 147, R. T. 272 (Aug. 1984), may be used to determine the susceptibility of a metal or alloy to localized corrosion. The RCP technique measures rupture or pitting potential (E_{r}) , repassivation potential (E_{p}) and current density (I_a) for a given metal or alloy. E_r is the potential at which the passivating film of a given material breaks down, and is directly related to the pitting tendency of the material in a particular environment. E_{p} is the potential at which active corrosion regions of the material are repassivated in a given environment. I_{a} is the current density which is a measure for the general corrosion rate. The higher Ia, the higher the corrosion rate. $E_{\rm r},\ E_{\rm p}$ and $I_{\rm a}$ are measured with a silver reference electrode and a working electrode constructed from the material subject to corrosive attack. The higher (more positive) the $\boldsymbol{E}_{\boldsymbol{r}}$ value, the more effective a given antifreeze formulation is in preventing pitting corrosion initiation and Similarly, a higher (more positive) $E_{\scriptscriptstyle D}$ value progress. indicates that the particular corrosion inhibitor formulation has a greater ability to repassivate existing pits and crevices. On the other hand the higher I, value the faster corrosion is proceeding, the less effective a particular corrosion inhibitor is for protecting a certain metal or alloy.

The RCP test procedure may be described as follows; polished specimens of the metal to be tested (working electrode) are immersed in a 30 percent by volume solution of a given antifreeze concentrate formulation in hard ASTM corrosive

5

10

15

20

25

15

20

25

water; that is water containing 148 mg/l sodium sulfate, 165 mg/l sodium chloride, 138 mg/l sodium bicarbonate (ASTM 1384 corrosive water) and in addition 364 mg/l calcium chloride mono hydrate.

Polarization is achieved by polarizing at a scan rate of 2 mV/second until rupture potential E_r is attained. A rapid increase in polarization current results at E_r as the protective passivating film breaks down. When the current reaches a predetermined maximum value, the scanning direction is reversed towards more cathodic potentials. The repassivation potential E_p is determined during this final phase of the RCP scan.

Table 2 shows the results of RCP measurements using the compositions of the invention on magnesium, aluminium and copper; measuring E_r , E_p and I_a . Adequate reduced corrosion rates are found for the invention examples (1 to 6).

The ASTM D4340 test is a standard test method which covers a laboratory screening procedure for evaluating the effectiveness of engine coolants in combatting corrosion of aluminium casting alloys under heat-transfer conditions that may be present in aluminium cylinder head engines. For the purposes of the present application, magnesium alloy was used instead of aluminium casting alloys. A heat flux is established through a cast magnesium alloy typical of that used in engine cylinder heads whilst exposed to an engine coolant under a pressure of 193 kPa. The temperature of the magnesium specimen is maintained at 135°C and the test is continued for one week. The effectiveness of the coolant for preventing corrosion of

WO 00/22189

10

15

20

the magnesium under heat-transfer conditions is evaluated on the basis of the weight change of the test specimen and also on the visual appearance. A heat-transfer corrosion cell is assembled according to the test methodology. The magnesium specimen is 6.5 cm in diameter, and 1.3 cm thick. The system is heated through a thermocouple inserted into a thermocouple hole in the test specimen. The test is carried out based upon 165 mg of radiant grade sodium chloride dissolved in 750 ml distilled or de-ionized water. 250 ml of the test coolant is then added. This amount is sufficient for two tests.

that corrosion protection at been found temperatures, as evaluated in the ASTM D4340 test on magnesium, is further significantly improved by the synergistic effect of preferably a fluoride and/or compound, fluor fluoro-carboxylate. Photographs of magnesium coupons are shown in Figure 1. Comparative example (C) shows very severe attack with deep pitting and deposits. Examples H, I and J show various gradations of minor attack and some deposits. The addition of a fluor compound in Examples 1 to 6, shows a dramatic improvement of corrosion protection and a significant reduction in deposit formation.

•					l able l	_				
		Monoacids		Diacids	ds	Alkylbenzoic acid	Hydro carbyl triazole	Silicate	Fluor Compound	punodi
Example	2-ЕНА	НА	νo	SA	DDA	PTBBA	TTZ	Si	HF	FC
¥	3%					•				
В							0.2%			
U									0.2%	
D		3%								
ய			3%							
í.					3%					
Ð				3%						
H			3%				0.2%		·	
Ь						3%	0.2%			
ſ			1.5%			1.5%	0.2%			
×			1.5%			1.5%	0.02%			
J			1.5%	,		1.5%				
W	0.5%		1.5%			1.5%	0.5%			
z			1.5%			1.5%	0.2%	0.1%		
_			1.5%			1.5%	0.2%		0.2%	
2						3%			0.2%	
3			3%						0.2%	
4			3%						0.2%	
\$						1.5%		-		1.5%
9			1.5%	0.2%		1.5%	0.2%		0.2%	

.

RCP measurements to determine corrosion inhibitor effectiveness (E, E, in mV, I, in $\mu A/cm^2$)

		_*	20	6.0	20	ю	٠	8	9	•			2	-	7	2			•		
	copper	ញ <u>ិ</u>	-10	09	100	01-			• .			1500	1200	1000	1040	1320					•
		យ័	830	640	120	006	•	920	006	•	-	1750	1600	1300	1320	1810	٠				
-		· l	35	25	20	50	70	30	40	30	20	20	15	15	20	20		01	01	30	20
	aluminium	យិ	•	-520	-750	460	•	•		580	-520	•	•	٠	•	>2400		-700	•		*
		ங்	-260	-400	-500	1390	<2400	096	340	1420	-340	>2400	>2400	>2400	>2400	>2400		300	1400	0 2 01	200
•		<u>.</u>	06	001	1000	001	001	. 001	001	40	35	. 02	30	30	30	30	20	30	20	001 .	30
•	magnesium	ញ្ច	•	•	•	•	•			-220		-1120	•	700	780	-20	-1000	•	700	•	• •
	_	យ៍	-1130	-1120	750	-240	<2400	.440	-1100	>2400	>2400	>2400	>2400	2200	>2400	2300	>2400	. 002	>2400	200	300
	example	•	∢	æ	ŭ	۵	ш	īr	Ö	Ξ	_	-,	'×	Σ	z		2	ю	4	\$	9

-20-

repassivation potential $\mathbf{E}_{\mathbf{p}}$ could not be determined

CLAIMS

15

20.

25

- 1. A corrosion-inhibitor formulation comprising
 - a) from 0.1 to 15 weight percent of one or more inhibitors selected from the group comprising an alkylbenzoic acid, a C_5 - C_{15} monobasic acid and a C_5 - C_{15} dibasic acid, or salts thereof; and
 - b) from 0.005 to 5 weight percent of a fluoride and/or a fluorocarboxylic acid, or salts thereof.
- 2. A corrosion-inhibitor formulation as claimed in Claim 1,
 wherein the alkylbenzoic acid is 4-tertbutylbenzoic acid.
 - 3. A corrosion-inhibitor formulation as claimed in Claim 1 or Claim 2, wherein said C_5 - C_{15} aliphatic monobasic acid is selected from the group consisting of octanoic acid, nonanoic acid, decanoic acid, undecanoic acid, dodecanoic acid, 2-ethylhexanoic acid and neodecanoic acid.
 - 4. A corrosion-inhibitor formulation as claimed in Claim 3, wherein said aliphatic monobasic acid is octanoic acid.
 - 5. A corrosion-inhibitor formulation as claimed in any preceding claim, wherein said C_5 - C_{15} aliphatic dibasic acid is selected from the group consisting of suberic acid, azelaic acid, sebacic acid, undecanedioic acid, -dodecanedioic acid, and the diacid of dicyclopentadiene.
 - 6. A corrosion-inhibitor formulation as claimed in any preceding claim further comprising a hydrocarbyl triazole or a thiazole.

- 7. A corrosion-inhibitor formulation as claimed in Claim 6, wherein said hydrocarbyl triazole or thiazole is present at from 0.005 to 1.0 weight percent.
- 8. A corrosion-inhibitor formulation as claimed in Claim 6 or Claim 7, wherein said hydrocarbyl triazole is tolyltriazole or benzotriazole.
 - 9. A corrosion-inhibitor formulation as claimed in any preceding claim, wherein the fluorocarboxylic acid is added as the alkali metal, ammonium or amine soluble salt of the acid.
- 10. A corrosion-inhibitor formulation as claimed in any preceding claim further comprising an additional corrosion inhibitor selected from the group comprising alkali metal borates, alkali metal silicates, alkali metal benzoates, alkali metal nitrates, alkali metal nitrites, and alkali metal molybdates.
 - 11. A corrosion-inhibitor formulation as claimed in any preceding claim for use with a heat transfer fluid comprising a liquid alcohol freezing point depressant.
- 12. A corrosion-inhibitor formulation as claimed in Claim20 11, wherein said liquid alcohol freezing point depressant is ethylene glycol.
 - 13. A corrosion-inhibiting coolant comprising a corrosion-inhibitor as claimed in any one of Claims 1 to 10, a liquid alcohol freezing point depressant, and water.

.10

15

20

- 14. A coolant as claimed in Claim 13, comprising from 25-75 volume percent water.
- 15. A heat transfer fluid formulation comprising:
 - o) from 65-99 weight percent of a water soluble liquid alcohol freezing point depressant;
 - p) from 0.1-15 weight percent of alkylbenzoic acid of general formula I

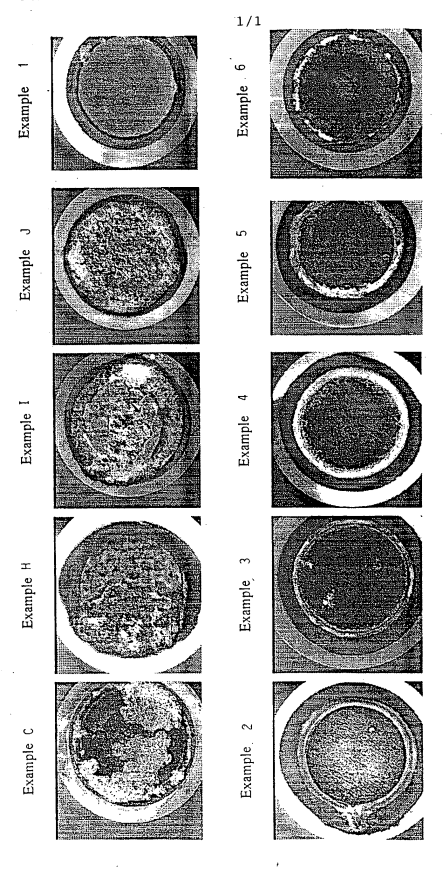
where R is a $C_1\text{-}C_\theta$ alkyl radical or an element of Group 7, or a salt thereof;

- q) from 0.1-15 weight percent of C_5-C_{15} aliphatic monobasic or dibasic acid or salt thereof;
- r) from 0.01-5 weight percent of hydrogen fluoride and/or a fluorocarboxylic acid or salt thereof.
- 16. The use of a corrosion-inhibitor formulation as claimed in any one of Claims 1 to 14 as a magnesium and/or aluminium corrosion-inhibitor in heat transfer fluids and engine coolants.
- 17. A process for inhibiting the general pitting, crevice and deposit-attack corrosion of magnesium and/or aluminium present in the cooling system of an internal combustion engine which comprises intermittently contacting the metal surface comprising magnesium and/or aluminium to be inhibited against corrosion with a heat transfer fluid comprising a

corrosion-inhibitor formulation as defined in any one of Claims.

1 to 10.

Figure 1: Pictures of test specimens after ASTM D4340 test



BEST AVAILABLE COPY

INTERNATIONAL SEARCH REPORT

Intern. ial Application No PCT/IB 99/01659

A CLASSIE	COATION OF SUBJECT MATTER		
IPC 7	C23F11/10 C23F11/18 C09K5/2	0	
	·		
	International Patent Classification (IPC) or to both national classifi	ication and IPC	
B. FIELDS S	cumentation searched (classification system followed by classification	ation symbols)	
IPC 7	CO9K C23F		·
		·	·_
Documentati	ion searched other than minimum documentation to the extent that	t such documents are included in the fields sea	arched -
	ata base consulted during the international search (name of data t	base and, where practical, search terms used)	
Flectronic da	ata base consulted during the linemational course, visite of		
	·		
	•		
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	· · · · · · · · · · · · · · · · · · ·	······································
Category °	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to daim No.
ļ			1.0
Α	EP 0 251 480 A (TEXACO PETROLEU	M)	1-8
	7 January 1988 (1988-01-07)		
	cited in the application claims; examples; table I		
A	EP 0 229 254 A (ASAHI GLASS CO	LTD ;ELNA	1
	CO LTD (JP)) 22 July 1987 (1987	-07-22)	:
1	cited in the application table 6		
			,
Α	DE 23 38 352 A (APPLIED CHEM PT	Y LTD)	· 1
	28 February 1974 (1974-02-28)		•
	claims 1,4		
		-/	
	,	·	
X Fur	rther documents are listed in the continuation of box C.	χ Patent family members are listed	I in annex.
° Special c	categories of cited documents :	"T" later document published after the int	emational filing date
"A" docum	nent defining the general state of the art which is not	or priority date and not in conflict with cited to understand the principle or the	Tine application out
cons	sidered to be of particular relevance r document but published on or after the international	invention	claimed invention
filing	date ment which may throw doubts on priority claim(s) or	cannot be considered novel or canno involve an inventive step when the d	ocument is taken alone
which	then twitch may throw bubbs of plonty carriers. It is cited to establish the publication date of another ion or other special reason (as specified)	"Y" document of particular relevance; the	claimed invention nventive step when the
"O" docur	ment reterring to an oral disclosure, use, exhibition or	document is combined with one or ments, such combination being obvious	nore other such docu-
"P" docum	r means ment published prior to the international filing date but	in the art. "&" document member of the same pater	
later	than the priority date claimed	Date of mailing of the international se	
Date of the	e actual completion of the international search	Date of maming of the michigational se	
	7 January 2000	17/01/2000	
<u> </u>		Authorized officer	
Name and	d mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2		
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni,	Puetz, C	
1	Fax: (+31-70) 340-3016	1 4002, 0	

PCT/IB 9	9/01659
----------	---------

ategory '	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
\ .	DATABASE WPI	1,2,5
`	Section Ch, Week 9613 Derwent Publications Ltd., London, GB;	
•	Class E19, AN 96-124252 XP002095342 & JP 08 020763 A (SEIKEN KAGAKU KOGYO KK),	
-	23 January 1996 (1996-01-23) cited in the application abstract	
1	PATENT ABSTRACTS OF JAPAN vol. 096, no. 008, 30 August 1996 (1996-08-30)	1,2,5
	& JP 08 085782 A (NIPPON CHEM KOGYO KK), 2 April 1996 (1996-04-02) cited in the application abstract	
A	EP 0 864 627 A (CCI CO LIMITED) 16 September 1998 (1998-09-16)	1,3
A	PATENT ABSTRACTS OF JAPAN vol. 016, no. 369 (C-0972), 10 August 1992 (1992-08-10) & JP 04 117481 A (SHII SHII AI KK), 17 April 1992 (1992-04-17) abstract	1,2,5,6
A	US 4 647 392 A (DARDEN JEROME W ET AL) 3 March 1987 (1987-03-03) cited in the application the whole document	1,3-6
Α	EP 0 457 859 B (SOLVAY INTEROX LIMITED) 27 November 1991 (1991-11-27)	1
		·
		,

INTERNATIONAL SEARCH REPORT

Information on patent family members

Intern. ial Application No PCT/IB 99/01659

Patent document cited in search report	t	Publication date	Patent family member(s)	Publication date
EP 0251480	A	07-01-1988	US 4851145 A	25-07-1989
	••		DE 3775634 A	13-02-1992
			JP 1753110 C	23-04-1993
•			JP 4042477 B	13-07-1992
			JP 63026388 A	03-02-1988
EP 0229254	 А	22-07-1987	JP 1707842 C	27-10-1992
		•	JP 3076007 B	04-12-1991
			JP 63050011 A	02-03-1988
			JP 1647938 C	13-03-1992
			JP 3006645 B	30-01-1991
			JP 62114208 A	26-05-1987
			US 4740869 A	26-04-1988
			US 4689724 A	25-08-1987
DE 2338352	Α	28-02-1974	AU 452099 B	29-08-1974
			AU 5798873 A	29-08-1974
			BE 803097 A	03-12-1973
		•	FR 2194802 A	01-03-1974
			GB 1434804 A	05-05-1976
			JP 49053545 A	24-05-1974
		·	NL 7310594 A	05-02-1974
JP 8020763	Α	23-01-1996	NONE	
JP 08085782	Α	02-04-1996	NONE	
EP 0864627	Α	16-09-1998	JP: 10251624 A	22-09-1998
JP 04117481	Α	17-04-1992	JP 2772578 B	02-07-1998
US 4647392	Α	03-03-1987	CA 1258162 A	08-08-198
			. EP 0229440 A	22-07-198
			JP 5073159 B	13-10-199
			JP 62158778 A	14-07-198
EP 457859	В	27-11-1991	AT 113670 T	15-11-199
			DE 69013896 D	08-12-199
			DE 69013896 T	09-03-199
			EP 0457859 A	27-11-199
			ES 2066226 T	01-03-199
			WO 9105079 A	18-04-199
		•		
			HK 1008058 A US 5364549 A	30-04-199 15-11-199